

Aqueous enzyme assisted oil extraction from oilseeds and emulsion deemulsifying methods: a review

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| 1 | Aqueous enzyme assisted oil extraction from oilseeds and emulsion de-emulsifying |
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| 2 | methods: a review |
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| 13 | |
| 14 | Abstract |
| 15 | Regulatory, safety, and environmental issues have prompted the development of aqueous |
| 16 | enzymatic extraction (AEE) for extracting components from oil-bearing materials. The |
| 17 | emulsion resulting from AEE requires de-emulsification to separate the oil; when enzymes |
| 18 | are used for this purpose, the method is known as aqueous enzymatic emulsion de- |
| 19 | emulsification (AEED). In general, enzyme assisted oil extraction is known to yield oil |
| 20 | having highly favourable characteristics. This review covers technological aspects of |
| 21 | enzyme assisted oil extraction, and explores the quality characteristics of the oils obtained, |
| | |

focusing particularly on recent efforts undertaken to improve process economics byrecovering and reusing enzymes.

24

25 Keywords

aqueous oil extraction, enzyme treatment, oil yield, oil characteristics, emulsion separation

28 **1. Introduction**

Aqueous enzymatic extraction (AEE) is a promising method for the simultaneous 29 extraction of oil and protein from oilseeds. The products are of superior quality and highly 30 31 suited to human consumption. In the extraction process, water containing selected enzymes 32 forms the extraction medium used for incubating the oilseeds. When enzymes are not employed, the process is termed as aqueous extraction which invariably results in lower oil 33 34 yield. The use of enzymes allows separation of targeted extracted components with unchanged properties which can potentially influence, favourably, the final product in 35 terms of taste and smell. Interest in this technological approach has also increased recently 36 37 due to safety and environmental regulatory concerns. In comparison with solvent extraction, the use of an aqueous medium is much safer, environmental-friendly and 38 economical. In addition, it contributes to a much safer and flexible operation, lower energy 39 40 consumption and operational costs, and lower capital investment. A variety of temporal crops can be processed, and the extracted oil does not need further refining. Non-toxic meal 41 42 and value-added fibre and protein are also produced as co-products, due to the milder operating conditions employed. In addition, the aqueous medium allows simultaneous 43

separation of phospholipids from the oil. Therefore, degumming step (in case of oilseeds)
is not necessary and the overall cost of processing can be reduced (Latif & Anwar, 2011;
Latif *et al.*, 2011; Yang Li *et al.*, 2011; Chabrand & Glatz, 2009; Jung & Mahfuz, 2009;
Wu *et al.*, 2009; Soto *et al.*, 2007; Santos & Ferrari, 2005; Gros *et al.*, 2003; Hanmoungjai *et al.*, 2001; Rosenthal *et al.*, 2001; Sineiro *et al.*, 1998; Ksenija *et al.*, 1997; Rosenthal *et al.*, 1996)

Despite the advantages, the application of AEE is still limited due to long 50 processing time and the high cost spent for the drying process after the enzyme treatment 51 52 (Shah et al., 2005; Dominguez et al., 1996). The high cost may also be attributed to the 53 enzymes themselves, because a significant amount is required (normally >1% of the weight of the oilseed taken). Further, the non-availability of enzymes on a commercial scale has 54 55 limited the development of such processes (Rui et al., 2009; Shah et al., 2005). An added 56 problem with AEE is that it is impossible to avoid emulsification of the extracted oil, which requires post extraction de-emulsification to recover and enhance oil yield (Latif & Anwar, 57 2011; Long et al., 2011; Wu et al., 2009; Chabrand et al., 2008; Santos & Ferrari, 2005; 58 59 Rosenthal et al., 1998; Sineiro et al., 1998a). Addition of suitable enzymes to the cream 60 emulsion may be able to separate the oil, and in this paper, this particular sequence of process is 61 termed as aqueous enzymatic emulsion de-emulsification (AEED). 62 In an earlier review by Rosenthal *et al.* (1996), the principles and mechanisms of:

mechanical, solvent, aqueous, and aqueous enzymatic extraction methods have been
addressed, besides reviewing the effects of enzymes on plant cell composition and methods
employed earlier for de-emulsification. The main purpose of this review is to critically
assess the information available to date, in order to conclude whether the enzymatic route

is a viable industrial option for any given oilseed. In addition, the other objectives of this
review are: to discuss the effect of incubating conditions in AEE on the oil extraction
efficiency; to compare AEE with other extraction methods in terms of yields and
characteristics of the oils from various oil-bearing materials; to explore methods available
to de-emulsify the oil- aqueous phase emulsions that are inevitably formed during
extraction; and finally, to explore the possibility of re-using in the enzyme after recovery in
order to make the process more cost effective.

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2. Aqueous enzymatic extraction (AEE) method Table 1 lists the enzymes used in 75 76 earlier research. In terms of the dispersion structure, Sineiro et al. (1998a) reported that 77 aqueous extraction resulted in oil droplets with spherical shapes in the case of sunflower oil. However, with the use of enzymes, the oil aggregates possessed different shapes with 78 79 less structured and irregular cell wall surface. Different oils exhibit different properties, and it is reasonable to assume that AEE of different oil-bearing materials result in oil droplets 80 81 with different characteristics. The enhancement in oil yield with the use of enzymes, i.e. 82 AEE as compared to aqueous extraction without enzymes from various oil-bearing materials are summarized in Table 2. The table also summarizes the differences observed 83 in oil yields between AEE and solvent extraction methods. It is clearly shown that the use 84 of enzymes increases the oil yield, yet it is still lower than the yield when solvent 85 86 extraction is used. Therefore, numerous studies have been conducted to establish the most 87 suitable enzymes that can be used, either individually or in combination, on various types 88 of oil-bearing materials in order to increase the oil yields.

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90 2.1. Studies comparing extraction efficiencies using different enzymes

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92 Figure 1(a) and 1(b) illustrate the flow sheets of AEE for soybean and olive oil, 93 respectively. The types of enzymes added depend on the cellular composition and structure 94 of the oil-bearing material (Passos et al., 2009). According to Rosenthal et al. (2001), the 95 use of Alcalase 2.4L (protease) increased the oil yield from heat-treated soybean flour as compared to cellulase, hemicellulase, and pectinase. Similarly, Santos and Ferrari (2005) 96 97 reported that both Alcalase and Celluclast (cellulase) were able to increase the oil yield 98 from soybeans, with Alcalase giving higher yields. A higher yield in the case of protease 99 (96.0%) as compared to phospholipase (73.4%) was also reported by Jung et al. (2009) in the case of extruded soybean flakes. In addition, Lamsal et al. (2006) reported that the use 100 101 of individual cellulase and a mixture of cellulase and protease did not significantly increase 102 the soybean oil yield from extruded soybean flakes (68%); yet the yield increased when 103 individual protease was added (88%). These findings illustrate the specificity of enzymes 104 and enzymatic mixtures for any given oil-bearing material. The presence of protein as a 105 major component in the cell wall of soybean seeds suggests that the oil is released more easily from the cellular matrix by degrading the proteins, which is achieved by the action of 106 107 protease. In the case of rapeseed, pectin is reported to be the major component of its cell 108 wall (Zhang et al. 2007), hence the highest oil yields, up to 85.9% in emulsified form, has 109 been reported when pectinase is used which is significantly greater than the values obtained 110 with other carbohydrases. Zhang et al. (2007) also employed a combination of pectinase

with cellulase and β -glucanase in a ratio of 4:1:1 to result in the highest yield (91.6% emulsified oil), this marginal enhancement in yield may be attributed to the elimination of other barriers to the release of oil. Similarly, Szydłowska-Czerniak *et al.* (2010) reported that the application of pectolytic enzyme (ROHAPECT PTE) under optimum conditions prior to pressing produced higher rapeseed free oil yield (16.5%) as compared to cellulolytic enzyme (15.5%).

Different from oilseeds, addition of enzymes is done on the olive paste in the case of olive fruits, followed by its kneading process as shown in Fig. 1(b). Most studies on extraction of olive oil involved addition of an enzyme mixture consisting mainly pectinase, cellulase, hemicellulase, and other minor enzymes. The studies also reported the inadequacies of these enzymes to extract olive oil if added individually (Aliakbarian et al., 2008; De Faveri et al., 2008; Chiacchierini et al., 2007).

In general, a better oil extraction yield can be expected when a judiciously chosen 123 124 mixture of enzymes is used because of possible synergy (Passos et al., 2009). However, 125 according to Rovaris et al. (2012), there was no significant difference in soybean oil yields when a mixture of Alcalase 2.4 L and Viscozyme was used as compared to a mixture of 126 127 Alcalase 2.4 L and Celluclast 1.5 L (29.48% as against 26.82% at pH 4.5; 20.63% as against 20.23% in the case of uncontrolled pH), even though Viscozyme itself is a mixture 128 of enzymes. There was also no significant difference in garlic oil yields upon addition of 129 130 Viscozyme as compared to addition of individual pectinase, protease, and cellulase as reported by Sowbhagya et al. (2009). A similar outcome was reported by Tabtabaei and 131 132 Diosady (2013) in yellow mustard flour oil extraction when Celluclast 1.5L and Pectinex Ultra SP-L were used, as against Viscozyme L. In addition, the use of Alcalase 2.4L and 133

134 Protex 7L resulted in highest sesame (Latif & Anwar, 2011) and Moringa oleifera (Latif et 135 al., 2011) seed oils, respectively, in comparison with Viscozyme L, Protex 7L, Natuzyme, 136 Kemzyme, and Multifect CX 13L which are essentially mixtures of enzymes (Latif 137 &Anwar, 2011; Latif et al., 2011). Viscozyme, being a mixture of enzymes, was reported 138 to have performed better in the case of sunflower oil extraction, which had been proved by Latif and Anwar (2009). A higher oil yield from bush mango kernel flour was also 139 observed upon addition of Viscozyme (68.0%) as compared to Alcalase (35.0%) and 140 141 Pectinex (42.2%) (Womeni et al., 2008). The different effects of the Viscozyme on oil 142 yields may be due to the nature of different oil-bearing materials and incubating conditions 143 employed.

In a different study conducted by Jiang et al. (2010), five different proteases were 144 145 tested to improve peanut oil yield, and the highest oil yield was obtained when Alcalase 146 was used (73.45%), followed by As1398 (66.36%), Nutrase (60.08%), Protizyme 147 (55.02%), and Protamex (48.89%). A combination of Alcalase with any of these enzymes 148 did not increase the oil yield. Therefore, Jiang et al. (2010) only used Alcalase which 149 reduced the extraction cost, and increased oil yield up to 79.32% under optimum incubating conditions. Similarly, the use of Neutrase 0.8L resulted in marginally lower 150 151 Moringa oleifera oil yield than when its combination with other three enzymes were 152 employed (Abdulkarim et al., 2006). In the case of flaxseed oil extraction conducted by 153 Long *et al.* (2011), the addition of cellulase, pectinase, and hemicellulase, individually, 154 gave higher yields than β -glucosidase and proteinase. Therefore, these authors used a 155 mixture of cellulase, pectinase, and hemicellulase (1:1:1) which resulted in a higher oil

yield of 61.7-66.1% as compared to the oil yield of each individual enzyme. With reference
to Table 2, , Zhang *et al.* (2007) reported highest yield of 92.7% in the case of rapeseed oil,
however, the oil remained very stably emulsified in the cream. Therefore, an alkaline
extraction was conducted by using Alcalase which resulted in protein degradation along
with an increase in total oil yield.

161 Based on the above studies, it is not possible to establish conclusively whether it is 162 better to use enzymes individually or in combination, although there are numerous 163 instances where there is a possibility that a mixture can work synergistically. The choice of 164 enzyme depends on the location of the oil within the cellular architecture and the 165 biochemical nature of the components surrounding it. It is therefore necessary, not only to 166 look at the dominant biochemical component holding the cellular matrix together, but also 167 investigate the cellular architecture and examine the specific components which act as a 168 barrier against the release of oil. It is only when both these factors are considered 169 simultaneously, the right enzyme mixture can be identified for a given oil-bearing material. 170

171 2.2. Studies on the use of enzyme as a pre-treatment step prior to extraction

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173 Recently, the application of enzyme pre-treatment prior to oil extraction has been 174 shown to increase yields (Li *et al.*, 2012). The addition of enzymes as a pre-treatment 175 weakens the cells and facilitate the following oil extraction methods such as mechanical 176 pressing and solvent treatment. Furthermore, the advantage of employing this approach lies 177 in the possibility of avoiding the formation of an oil-in-water emulsion that is very difficult

| 178 | to separate after the extraction processes. The reported enhancement in oil yields with the |
|-----|---|
| 179 | use of enzyme pre-treatment is summarized in Table 3. In addition to the higher yield, |
| 180 | Dominguez et al. (1996) also reported that it was easier to extract the sunflower oil |
| 181 | remaining in a mass of pre-treated mechanically pressed cake. In the case of Chilean |
| 182 | hazelnuts, enzyme pre-treatment resulted in significantly lower residual oil in the meal as |
| 183 | reported by Zuniga et al. (2003). Overall, these studies indicate that enzyme pre-treatment |
| 184 | is applicable to various oil-bearing materials and can be employed prior to both mechanical |
| 185 | and solvent extraction methods. The oil yield enhancement is due to the hydrolytic action |
| 186 | of the enzymes on the cell wall and membrane components which facilitate subsequent oil |
| 187 | release. |
| 188 | |
| 189 | 2.3. Studies on pre-treatment step prior to enzymatic extraction |
| 190 | |
| 191 | Some studies have highlighted potential pre-treatment methods, which are not |
| 192 | necessarily enzyme-based that could be followed up by AEE as summarized in Table 4. In |
| 193 | the case of high pressure processing as reported by Jung and Mahfuz (2009), the use of |
| 194 | high pressure induced protein aggregation yet it was further hydrolyzed by protease, thus |
| 195 | facilitated oil removal. On the other hand, Shan Liu et al. (2011) reported that ultrasound |
| 196 | generated cavitations which accelerated the leaching out of cellular components including |
| 197 | oil. The use of extrusion prior to AEE has been extensively studied by Jung and Mahfuz |
| | |

- 198 (2009), Jung *et al.* (2009), and Wu *et al.* (2009). According to these authors, protein
- aggregates are formed during extrusion but these entrap or interact with the oil. The

| 200 | interactions could then be disrupted by the use of protease, which result in increasing the |
|-----|--|
| 201 | oil and protein yields. These studies have shown the potential of AEE assisted by other pre- |
| 202 | treatment methods to increase oil yields. |
| 203 | |
| 204 | 2.4. Factors affecting the efficiency of enzymatic extraction |
| 205 | |
| 206 | Table 5 summarizes the maximum oil yields resulting from various oil-bearing |
| 207 | materials as influenced by the selected and optimized incubating conditions. The key |
| 208 | factors affecting the efficiency of AEE will be discussed separately, below. |
| 209 | |
| 210 | 2.4.1.Particle size of the oil-bearing materials |
| 211 | Most of the early studies did not consider the particle size of the oil-bearing |
| 212 | material as a key factor influencing extraction efficiency (Passos et al., 2009; Rosenthal et |
| 213 | al., 2001). Theoretically, the lower the particle size, the higher the oil yield for a given set |
| 214 | of extraction conditions, which is attributable to higher cell wall disruption during size |
| 215 | reduction as well as the lower diffusion path length for both enzymes and cellular |
| 216 | components. However, according to Passos et al. (2009), materials with high oil content |
| 217 | but exhibiting a weak structure, may collapse and lose their microporosity when treated |
| 218 | with solvents, which can result in non-uniform percolation and be detrimental to extraction |
| 219 | efficiency. In addition, grinding of materials with high oil content into very low particle |
| 220 | sizes may cause the particles to adhere, as reported by Nyam et al. (2009a) in the case of |
| 221 | Kalahari melon seeds. Therefore, in industry, starting materials with very low particle size |
| | |

222 are not recommended and there appears to be an optimum size. This illustrates the 223 importance of selecting the right particle size prior to extraction as had been done by some authors. Sineiro et al. (1998a) used ground soybean and sunflower seeds having mean 224 225 particle size <0.2 mm. The grape seeds used by Passos et al. (2009) were grouped into 226 different particle size ranges (in mm): <0.50, 0.50-0.60, 0.60-0.71, 0.71-1.0, 1.0-1.4, 1.4-227 2.0, and >2.0, and increment in oil yield was observed at lower particle sizes. In the case of linseed oil, Gros et al. (2003) reported no oil recovery from whole linseed kernels, because 228 229 the substrate was not accessible to the enzymes added. Instead, the hull broke down and the 230 kernels expanded due to hydration. On the other hand, when the kernels were crushed to 231 form different particle sizes including fine powders, the yields improved, particularly after 232 applying hydraulic pressures (Gros et al., 2003). Similarly, in the case of soybean, the use of flour resulted in 24% higher yield than the flakes (Jung et al., 2009), while 31% yield 233 234 enhancement was reported by Rosenthal et al. (1998) when the particle size was reduced from 400 μ m to 100 μ m. 235

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237 2.4.2. Enzyme/substrate ratio

Higher enzyme concentration leads to greater interaction between the enzyme and
substrate, thus promoting cell wall degradation and rupturing more peptide bonds (Teixeira *et al.*, 2013; Jiang *et al.*, 2010; Dominguez *et al.*, 1996). However, too high enzyme
concentration may result in bitterness and off flavours, as reported by Jiang *et al.* (2010),
possibly due to the extraction of undesirable components. Most authors have reported
similar trends where the oil yield increased up to certain enzyme concentration only,

followed by steady or decreased rate which may be due to saturation of the substrates

245 (Jiang *et al.*, 2010), or caramelization of soluble sugars that limit oil release (Zuniga *et al.*,

246 2003). In general, the actual concentration used will depend on process economics

especially the cost of enzymes (Long *et al.*, 2011; Zhang *et al.*, 2007), and the quality of

the oil extracted.

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250 2.4.3. Ratio of water to oil-bearing material

251 The water used in AEE not only serves as an extraction medium but also enters the 252 oil-bearing material and modifies its water activity. The resulting moisture content of the 253 oil-bearing material can assist hydrolytic reaction, diffusion, and mobility of the enzymes 254 and products (Yang Li et al., 2011; Zhang et al. 2007; Sineiro et al., 1998a; Dominguez et 255 al., 1996). On the other hand, very low moisture content results in the formation of thick 256 suspensions which can prevent the enzymes from effectively penetrating into the substrate (Zhang et al., 2007). Sineiro et al. (1998a) reported that only certain 'areas' in sunflower 257 kernels were degraded by enzymes at low moisture content. Although, materials with 258 259 higher water activity demonstrate higher extraction efficiency (Soto *et al.*, 2007), the 260 presence of excessive moisture content in the oil-bearing material can decrease the 261 concentration of enzymes and substrates, and have an adverse effect on extraction (Yang Li 262 et al., 2011; Zhang et al., 2007; Dominguez et al., 1996). Therefore, selection of 263 appropriate moisture content is critical for the success of AEE. 264

265 2.4.4. pH of extraction medium

266 The pH at which enzymes attain maximum activity varies with the enzyme. In most 267 earlier studies, the pH value of the solution, be it for soaking pre-treatment or extraction 268 itself, was set at a value corresponding to maximum enzyme activity (Latif & Anwar, 2011; 269 Jung & Mahfuz, 2009; Wu et al., 2009; Abdulkarim et al., 2005; Rosenthal et al., 2001; 270 Sineiro *et al.*, 1998). However, the optimum pH of a number of enzymes is in the range of 271 the isoelectric pH of proteins which depends on the nature of the oilseeds; since proteins 272 are highly insoluble in this range of pH, oil release may get inhibited. Therefore, the pH 273 value employed must not only be conducive for the action of enzymes but it should also be remote from protein isoelectric point (Tabtabaei & Diosady, 2013; Wu et al., 2009; Sineiro 274 275 et al., 1998; Rosenthal et al., 1996). This is yet another reason why many authors 276 considered using a mixture of enzymes which demonstrates high activity at pH values 277 remote from the isoelectric point and remain effective for oil extraction. The enzymes are 278 able to solubilize and hydrolyze the proteins besides disrupting other polysaccharide 279 constituents which facilitate oil release (Rovaris et al., 2012; Latif & Anwar, 2011; Passos et al., 2009). Long et al. (2011) had used a mixture of cellulase, pectinase, and 280 281 hemicellulase (1:1:1) at pH 4.5-5.0 which resulted in highest flaxseed oil yield (73.9%) as compared to oil yield of each individual enzyme. In the case of soybean oil, at pH 4.5, 282 Rovaris et al. (2012) used a mixture of Alcalase 2.4L and Celluclast 1.5L which resulted in 283 284 26.82% oil (20.63% in the case of uncontrolled pH), and a mixture of Alcalase 2.4 L and Viscozyme which resulted in 29.48% oil (20.23% in the case of uncontrolled pH). A 285 number of studies have also used ProtizymeTM for the AEE (Jiang et al., 2010; Gaur et al., 286 2010; Sharma *et al.*, 2002). ProtizymeTM, being a mixture of proteases, possess different 287

optimum pH which allowed selection of any incubating pH sensitive to the isoelectric point
of the major protein fraction of the seeds. Overall, proper pH selection critically influences
yields of oil and other components in AEE .

291

292 2.4.5. Incubation temperature

293 Besides being active over a narrow range of pH, enzymes also active over a narrow 294 temperature interval. According to Rui et al. (2009), the optimum temperature range for enzymatic hydrolysis is between 40-55 °C, thus many authors employ AEE temperatures 295 296 which fall within this range. In practice, one often prefers to use the lowest possible 297 temperature yielding adequate activity (Passos *et al.*, 2009). In the case of olive fruits, a lower temperature of 30 °C was found to be favourable especially to preserve the oil 298 299 quality (Aliakbarian et al., 2008; De Faveri et al., 2008; Ranalli et al., 2003; Garcia et al., 300 2001; Ranalli et al., 1999). Gros et al. (2003) also used a temperature of 34 °C for similar 301 reason in linseed oil extraction. A significant effect of temperature on oil yield was 302 reported by Sharma et al. (2002), where highest peanut oil yield was observed at 40 °C, but 303 it decreased significantly when the temperature was reduced to 37 °C. According to Zúniga et al. (2003), at temperatures greater than 45 °C, enzymatic hydrolysis begins to decrease 304 305 due to enzyme inactivation which leads to lower oil yield. The oil release from the cells 306 may also be limited due to presence of soluble sugars in the composition which can 307 undergo caramelization during the drying stage. Therefore, similar trends were reported 308 from most of the conducted studies, where the oil yield increased up to certain temperature 309 only, followed by steady or decreased rate afterwards. Thus, besides the oil yield, the oil

310 quality characteristics must also be taken into consideration when selecting AEE

311 temperature.

312

313 2.4.6. Incubation time

314 According to Jiang et al. (2010), Abdulkarim et al. (2006), Santos and Ferrari 315 (2005), and Dominguez et al. (1996), degradation of cell wall components can be enhanced 316 by prolonging the incubation time. Passos et al. (2009) also reported that the use of an 317 enzyme mixture of cellulase, protease, xylanase, and pectinase for 120 hr resulted in 3.8% 318 higher yield as compared to 24 hr of incubation time. However, this time duration (i.e. 120 319 hr) is far too long to be acceptable in practice (Passos et al., 2009), lower oil quality may 320 result (Jiang et al., 2010), leading to high energy usage and production of undesirable 321 products (Abdulkarim et al., 2006). In addition, Rui et al. (2009) highlighted that longer 322 incubation time of AEE in relation to other solvent extraction methods is one of the 323 disadvantages of AEE. In some cases, the oil yield decreased after a certain incubation 324 period because the whole substrates have reacted with the enzymes; leaving negligible 325 substrates left for further enzymatic reaction to take place (Zhang et al., 2007). On the 326 whole, these studies have shown that although oil yield may increases with time, the rate of 327 increase may be far too slow to warrant extended operations, and the oil quality may also 328 get compromised.

329

330 2.4.7. Agitation rate

| 331 | According to Rosenthal et al. (1998) and Sineiro et al. (1998a), agitation assists in |
|-----|--|
| 332 | mixing and additional rupture of the cell wall, and agitation rate is one of the factors |
| 333 | affecting the disruption of cell wall. Abdulkarim et al. (2006) reported that the agitation |
| 334 | rates of 50 and 80 rpm were not adequate to separate the Moringa oleifera oil from other |
| 335 | seed components, thus resulted in lower oil yield than at 120 rpm. At this agitation rate of |
| 336 | 120 rpm, bigger oil droplets were observed to accumulate at the surface which enabled |
| 337 | easier separation. A similar observation was reported at 80 rpm in extraction of peanut oil |
| 338 | (Sharma et al., 2002) and at 100 rpm in the extraction of Kalahari melon seed oil (Nyam et |
| 339 | al., 2009a). On the other hand, the use of higher speeds leads to higher energy consumption |
| 340 | and cost (Rosenthal et al., 1998), besides resulting in the formation of a more stable oil- |
| 341 | aqueous phase emulsion that is difficult to separate (Nyam et al., 2009a; Abdulkarim et al., |
| 342 | 2006; Sharma et al., 2002, Hanmoungjai et al., 2000). These studies highlight the |
| 343 | importance of selecting appropriate agitation rate that will result in the highest oil yield |
| 344 | possible, considering both the oil recovered and emulsion stability at the end of the AEE |
| 345 | process. |
| 346 | |
| 347 | 2.5. Multi factorial studies on AEE |
| 348 | |
| 349 | A number of authors have employed statistical methods to indicate the relative |
| 350 | importance of the AEE parameters listed above. According to Rosenthal et al. (2001), |

352 of the ground seeds, the ratio of water to oil-bearing material, and the interaction between

351

soybean oil yield was significantly influenced by the type of enzyme used, the particle size

353 the two latter parameters. However, according to Hanmoungjai et al. (2001), only the 354 enzyme concentration had the most significant effect on the extraction of rice bran oil, while both the incubation time and temperature did not significantly affect the oil yield. 355 356 Different AEE parameters used for other samples such as bayberry kernels (Zhang et al., 357 2012), kalahari melon seeds (Nyam et al., 2009a), palm fruit (Teixeira et al., 2013), peanuts 358 (Jiang et al., 2010), and pine kernels (Yang Li et al., 2011) also had different degree of 359 significant effect on oil yield. These studies show that it is almost impossible to generalize which factor is important and which is not, for a given material. It is necessary to undertake 360 361 an experimental investigation before designing and scaling up an AEE process.

362

363 3. De-emulsification methods for aqueous enzymatic process (AEED)

364 When oil is extracted into an aqueous enzymatic phase, it inevitably forms an emulsion, 365 which is often difficult to separate because of the added stability imparted by the 366 interfacially active cellular components which are also extracted in the same process. It is 367 therefore necessary to carefully consider the techniques employed to separate the oil, 368 because the final yield and oil quality, and the economic viability of the process, will depend critically on de-emulsification steps. When AEE is followed by a centrifugation 369 370 step, besides oil, other fractions recovered include a skim and a cream emulsion (Figure 371 1(a)). The cream emulsion is very stable due to its protein content which acts as an 372 excellent emulsifier. Addition of suitable enzymes to the cream emulsion may be able to 373 separate the oil, and in this paper as had been mentioned earlier, this particular sequence of 374 process is termed as aqueous enzymatic emulsion de-emulsification (AEED). The enzymes 375 used in the AEED processes were also listed in Table 1. In this method, the enzymes added 376 to the cream emulsion hydrolyze the interfacial proteins, thus reducing their molecular size and decreasing the rigidity of the oil droplet interface. The enzymes also remove the high 377 378 molecular weight polypeptides which may occupy the emulsion interface and further 379 reduce the interfacial membrane thickness. These enzymatic reactions lead to greater oil 380 droplet coalescence and assist in free oil release (Tabtabaei & Diosady, 2013; Raghavendra 381 & Raghavarao, 2010; Chabrand & Glatz, 2009; Jung & Mahfuz, 2009; Marina et al., 2009; Wu et al., 2009; Chabrand et al., 2008). The original enzymes used in the AEE may also be 382 383 carried out into the cream emulsion and assist hydrolytic reactions if suitable incubating 384 conditions were employed (Chabrand & Glatz, 2009; Jung et al., 2009). The free oil yield is commonly expressed as a percentage based on the initial weight of the cream emulsion. 385 In the case of oil-bearing coconut milk, the emulsion needs to be destabilized in 386 387 order to obtain virgin coconut oil as shown in Figure 1(c). According to Jena and Das (2006), Garcia et al. (2005), Tangsuphoom and Coupland (2005), and Balasundaresan et al. 388 389 (2002), coconut milk emulsion is low in stability due to its high fat content and the 390 presence of coconut proteins (~65% is globulin known as cocosin) with low emulsifying properties. Therefore, these authors noted that the separation was not too challenging and 391 392 concluded that the oil droplets were prone to undergo aggregation and tended to separate. 393 In contrast, Marina et al. (2009), Tangsuphoom and Coupland (2008), Peamprasart and 394 Chiewchan (2006), and McGlone *et al.* (1986) reported that a coconut cream emulsion was 395 highly stable due to presence of natural phospholipids and coconut proteins (mainly 396 globulins and albumins) which requires extra energy to be destabilized. It is not uncommon

| 397 | to find such conflicting reports in literature, in this area, which is principally because, most |
|---|---|
| 398 | papers do not take a holistic view on the whole process. Whether the downstream de- |
| 399 | emulsification is challenging or not depends on the process conditions employed during |
| 400 | AEE. If the conditions employed are such that the emulsion formed is very stable, then the |
| 401 | de-emulsification will naturally become challenging. On the other hand, careful process |
| 402 | design upstream, and use of conditions that do not favour the formation of a stable |
| 403 | emulsion whilst releasing significant yields of oil, will simplify de-emulsification and |
| 404 | enhance free oil yields and oil quality. |
| 405 | |
| 406 | 3.1. Studies comparing different enzymes for de-emulsification of cream emulsion |
| 407 | |
| | |
| 408 | Table 5 summarizes the types of enzymes and the incubating conditions used in |
| 408 409 | Table 5 summarizes the types of enzymes and the incubating conditions used in AEED methods for maximum free oil yields. In the case of yellow mustard flour, Tabtabaei |
| | |
| 409 | AEED methods for maximum free oil yields. In the case of yellow mustard flour, Tabtabaei |
| 409 410 | AEED methods for maximum free oil yields. In the case of yellow mustard flour, Tabtabaei and Diosady (2013) reported that Protex 6L possessed greater efficiency in the de- |
| 409 410 411 | AEED methods for maximum free oil yields. In the case of yellow mustard flour, Tabtabaei and Diosady (2013) reported that Protex 6L possessed greater efficiency in the de- emulsification process, as compared to other proteases and carbohydrases tested. Lipomode |
| 409 410 411 412 | AEED methods for maximum free oil yields. In the case of yellow mustard flour, Tabtabaei and Diosady (2013) reported that Protex 6L possessed greater efficiency in the de- emulsification process, as compared to other proteases and carbohydrases tested. Lipomode (Phospholipase A2), being one of the carbohydrases, resulted in the production of |
| 409 410 411 412 413 | AEED methods for maximum free oil yields. In the case of yellow mustard flour, Tabtabaei and Diosady (2013) reported that Protex 6L possessed greater efficiency in the de- emulsification process, as compared to other proteases and carbohydrases tested. Lipomode (Phospholipase A2), being one of the carbohydrases, resulted in the production of lysophospholipids which is an emulsifier, thus increased the emulsion stability and |
| 409 410 411 412 413 414 | AEED methods for maximum free oil yields. In the case of yellow mustard flour, Tabtabaei and Diosady (2013) reported that Protex 6L possessed greater efficiency in the de- emulsification process, as compared to other proteases and carbohydrases tested. Lipomode (Phospholipase A2), being one of the carbohydrases, resulted in the production of lysophospholipids which is an emulsifier, thus increased the emulsion stability and decreased the free oil yield. Lysophospholipids also present in small amount in G-ZYME |
| 409 410 411 412 413 414 415 | AEED methods for maximum free oil yields. In the case of yellow mustard flour, Tabtabaei and Diosady (2013) reported that Protex 6L possessed greater efficiency in the de- emulsification process, as compared to other proteases and carbohydrases tested. Lipomode (Phospholipase A2), being one of the carbohydrases, resulted in the production of lysophospholipids which is an emulsifier, thus increased the emulsion stability and decreased the free oil yield. Lysophospholipids also present in small amount in G-ZYME G999, resulted in an insignificant increase in the free oil yield. In the case of soybean oil, |

| 419 | have also reported that the use of enzymes shown in Table 5 at their optimum pH and |
|-----|---|
| 420 | temperature resulted in total de-emulsification of the cream emulsions, either the enzymes |
| 421 | had been used individually or in combination, or sequentially. These studies indicated that |
| 422 | the free oil yield depends on the stability of the cream emulsion which is mainly affected |
| 423 | by the AEE, besides the incubating conditions of the AEED which are discussed below. |
| 424 | |
| 425 | 3.2. Factors affecting the efficiency of enzymatic de-emulsification |
| 426 | |
| 427 | 3.2.1. Enzyme concentration |
| 428 | Generally, the use of higher enzyme concentration resulted in higher free oil yield. |
| 429 | According to Jung et al. (2009), at 25 °C, the use of Protex 6L resulted in higher free |
| 430 | soybean oil yield of 96% at 2.5% (w/w) concentration when compared to a 85-89% yield |
| 431 | while employing enzyme at 1.25% (w/w). Similarly, Wu et al. (2009) reported that free |
| 432 | soybean oil yield increased with increasing enzyme concentration starting from 0.2% |
| 433 | (w/w). In this study, when the LysoMax ^{TM} enzyme was used at a concentration lower than |
| 434 | 0.2% (w/w), the enzyme modified soybean phospholipids and caused the production of an |
| 435 | emulsifier known as lysolecithin. This emulsifier enhanced the stability of the cream |
| 436 | emulsion and therefore resulted in lower free oil yield. In addition, according to Wu et al. |
| 437 | (2009), increasing the LysoMax TM enzyme concentration did not increase the oil droplets |
| 438 | size. These authors also reported that in the concentration range of 0.2-2.0% (w/w), the use |
| 439 | of Protex 51FP resulted in higher free oil yield as compared to the LysoMax TM which |
| 440 | indicated the dominant role of soybean protein in stabilizing the cream emulsion. |

441

442 *3.2.2. pH value*

As had been discussed earlier (section 2.4.4), different enzymes possess different 443 444 optimum pH where maximum activity is observed. Therefore, most studies employed the optimum pH of the enzyme used in order to obtain the highest free oil yield (Table 5). In 445 the case of soybean oil, according to Wu et al. (2009), the oil droplet size and free oil yield 446 increased when the pH was lowered to 4.5, but not lower than 4.0. At the pH of 4.5, which 447 is the isoelectric point of soy protein, electrostatic repulsion between oil droplets decrease, 448 449 thus further enhancing oil droplets coalescence, formation of larger oil droplets, and higher 450 free oil yield (Wu et al., 2009). In a study conducted by Chabrand and Glatz (2009), the authors reported as high as 83% free soybean oil yield when the pH of the cream emulsion 451 452 was reduced to pH 4.5, and addition of enzyme (G-ZYME G999) at this similar pH 453 increased the free oil yield up to 100%. Similarly, Wu et al. (2009) reported that the use of G-ZYME G999 and Protex 50FP separately at pH 4.5 resulted in 100% free oil yield. 454 These authors suggested that the combination of enzymatic reaction and pH reduction leads 455 456 to coalescence of the oil droplets and formation of much bigger droplets than when enzymes are not used. Chabrand and Glatz (2009) had also reported the use of high pH on 457 the free soybean oil yield. At pH 9, only 2% of free oil yield was recovered. With the use 458 459 of enzymes (i.e. AEED) at pH 8 which was the original pH of the cream emulsion, no free 460 oil yield was obtained. Similarly, Wu et al. (2009) reported that the free soybean oil yield 461 decreased when the pH was increased beyond pH 4.5 up to pH 8. Therefore, the 462 significance of enzymes addition at suitable pH values for higher free oil yield is clear.

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3.2.3. Incubation time and temperature

| 465 | Similar to the pH value, different enzymes possess different optimum temperature |
|-----|---|
| 466 | where maximum activity is observed. Therefore, most earlier studies employed the |
| 467 | optimum temperature reported for the enzyme used in order to obtain highest free oil yield |
| 468 | (Table 5). Jung et al. (2009) reported the effect of different de-emulsification temperatures |
| 469 | and times on the free soybean oil yield when Protex 6L was used. Prolonged incubation |
| 470 | time from 2 min to 90 min enhanced the free oil yield from 86% to 100% at 65 °C. |
| 471 | However, the incubation time did not affect the free oil yield at lower temperatures of 25 |
| 472 | °C and 50 °C. Increment of temperature from 50 °C to 65 °C also increased the free oil |
| 473 | yield from 90% to 100% after incubation for 90 min. In the case of coconut milk de- |
| 474 | emulsification, Raghavendra and Raghavarao (2010) reported a higher free oil yield when |
| 475 | the use of enzyme was followed by chilling and thawing. In this case, a higher free oil yield |
| 476 | of 94.5% was reported at a higher temperature of 37 °C as compared to 91.0% yield at 25 |
| 477 | °C, because according to these authors, most enzymes possess an optimum temperature of |
| 478 | 37 °C. In addition, chilling resulted in packed oil bodies which are easier to separate |
| 479 | (Raghavendra & Raghavarao, 2010). |
| 480 | It is also possible to demulsify without the use of enzymes as reported by Jung et al. |
| 481 | (2009). In this study, the increase in temperature from 50 °C to 65 °C increased the free oil |
| | |

yield from 75% to 94%. According to the authors, the significant increase in free oil yield

may be due to the action of remaining protease in the cream emulsion which was carried 483

out from the AEE. In the case of yellow mustard flour, Tabtabaei and Diosady (2013) 484

subjected the emulsion recovered after AEED process to an alkaline treatment whichresulted in higher oil yield than AEED alone.

487 Other processing parameters such as shaking, de-canting, and stirring may also
488 influence de-emulsification efficiency (Jung *et al.*, 2009).

489

490 **4. Oil characteristics**

491 Most authors have reported the effects of extraction methods on the oil characteristics which are summarized in Table 6. With reference to the table, the oil yields from most of 492 493 the enzyme treatments were lower in oxidative deterioration and rancidity, indicated by the 494 lower free fatty acids and peroxide values as compared to the yields from solvent 495 treatments. It was assumed that the high temperature used during the solvent extraction 496 resulted in lower oxidative quality of the oils (Latif et al., 2011; Latif & Anwar, 2011; Latif 497 & Anwar, 2009; Latif et al., 2008). The peroxide value of rice bran oil extracted by solvent was also higher than that extracted enzymatically, but the difference was too small to the 498 limit industrial application (Hanmoungiai et al., 2001). In contrast, Kalahari melon seed oil 499 500 from AEE process gave higher free fatty acid and peroxide value than solvent extracted oil. 501 This may be due to the lipase activity in the seeds during the initial heating in the case of 502 AEE process (Nyam et al., 2009).

With reference to Table 6, some of the enzymatically extracted oils gave higher iodine value (IV) than aqueous and solvent extracted oils. Hanmoungjai *et al.* (2001) and Long *et al.* (2011) reported that the higher IV indicated higher polyunsaturated fatty acid content which therefore suggested a higher antioxidant activity. In addition, highest total

507 tocopherols was observed in most seed oils obtained from the AEE, followed by aqueous 508 and solvent extracted oils. It was suggested that the higher temperature employed in the 509 solvent treatment reduced the tocopherol content in the oil (Latif et al., 2011; Latif & 510 Anwar, 2011). The total tocopherols in olive oils reported by Ranalli et al. (2001) and 511 Ranalli et al. (2003) were also higher when AEE was employed as compared to aqueous 512 extractions without enzymes. In contrast, Nyam et al. (2009) reported lower total 513 tocopherol content in the Kalahari melon oil obtained by AEE than solvent extraction 514 method. This may be due to the production of components during the digestion process in 515 the AEE that can influence the amount of non-saponifiable matter, including tocopherols 516 (Gunstone, 2000),

In terms of total phenolic content, the values varied with different oil-bearing 517 materials, extraction methods employed, and the types of enzymes used in the AEE 518 519 process. In the case of olive oil, AEE resulted in higher total phenolic content than the aqueous extractions without enzymes. This may be due to cell wall hydrolysis by the 520 521 enzymes used which further assists partitioning of the phenolics into the oil. The phenolic 522 content positively influences oxidative stability, shelf life, nutritional, sensory, and health properties of the olive oil, besides flavour which got a greater sensory score (Latif & 523 Anwar, 2009, 2011; Aliakbarian et al., 2008; Ranalli et al., 2003; Ranalli et al., 1999; 524 525 Ranalli & De Mattia, 1997). Najafian et al. (2009) also reported that at higher enzyme 526 concentration, the phenolic content increased whilst the oil turbidity decreased, which may 527 be due to the enzymatic effect in reducing the amount of colloidal particles.

528 In terms of the fatty acid compositions (FAC), most authors reported similarities 529 between the oils obtained from solvent and enzymatic extraction methods (Teixeira et al., 2013; Li et al., 2012; Zhang et al., 2012; Latif et al., 2011; Latif & Anwar, 2009, 2011; 530 531 Jung et al., 2009; Nyam et al., 2009, 2009a; Latif et al., 2008). In a study conducted by Rui 532 et al. (2009), the FAC of the pitaya oil obtained from microwave-pre-treated enzyme 533 treatment was similar to the recommended FAC by the US dietary standard. Rui et al. (2009) suggested that microwave irradiation enhanced volumetric swelling of the cells in 534 the seed kernels which caused cell walls rupture, while the enzymes hydrolyzed the cell 535 536 wall and the bonds between the protein or pectin. A combination of these methods led to 537 extraction of pitaya oil with varying fatty acid types as compared to other methods. In the case of flaxseed oil, Long et al. (2011) reported that the oil yield from enzyme-pre-treated 538 ultrasonication possessed higher monounsaturated and polyunsaturated fatty acids than the 539 540 flaxseed oil obtained by solvent extraction. According to the authors, the use of water allowed diffusion of water-soluble components instead of the oil. Therefore, the oil 541 542 possessed approximately similar FAC as the original flaxseed oil (Long *et al.*, 2011). 543 In addition to the characteristics listed in Table 6, the colour intensity of oil had also been reported in some studies based on red and yellow units; higher values of these units 544 correspond to higher colour intensity. In the case of *Moringa oleifera* seeds, according to 545 546 Latif et al. (2011) and Abdulkarim et al. (2006), the different enzymes used in the AEE processes act on different components of the seeds which resulted in oil yields having 547 548 different colour intensity. However, the difference was more significant between the oil obtained by AEE and solvent extraction methods, which is similar to the results reported by 549

550 Nyam *et al.* (2009) and Latif *et al.* (2008) for Kalahari melon and canola seed oil,

respectively. The solvent-extracted oil had higher colour intensity which may due to the
pigments extracted by the solvent into the oil, such as carotenes and chlorophylls. The oil
obtained from AEE process may not need refining due to low colour intensity which
reduces the processing costs (Latif & Anwar, 2009; Nyam *et al.*, 2009; Latif *et al.*, 2008;
Abdulkarim *et al.*, 2006, Abdulkarim *et al.*, 2005).

Besides the colour of the oils, the sterols were also significantly lower in oil 556 557 obtained by AEE than solvent extracted oil, which suggests the ability of the solvent used 558 to extract lipid-soluble components (Nyam et al., 2009). In addition to these characteristics, 559 Sowbhagya et al. (2009) reported that the use of enzymes as a pre-treatment prior to steam distillation or hydrodistillation resulted in garlic oil with higher concentration of dithiins 560 561 which possess health benefits and highly desirable from a nutraceutical point of view. In 562 the case of soybean oil, with the use of enzymes, Jung et al. (2009) reported lower phosphorus content (<200ppm) which comply with the specification of the National 563 564 Oilseed Processors Association trading rules for crude degummed soybean oil. In a study 565 done by Ranalli et al. (1999), the Cytolase 0 enzyme used in olive oil extraction was harmless and water-soluble. Therefore, after the enzyme exerted all its effects on oil 566 extraction, it came out into the water (i.e. olive juice) and left no residue in the oil. Thus the 567 568 olive oil composition was not modified.

In extraction of virgin coconut oil from coconut milk emulsion, a combination of
AEED, chilling, and thawing for the coconut milk destabilization resulted in highest
creaming index as compared to other destabilization methods which indicated faster oil

| 572 | droplets movement and higher droplets aggregation. As compared to commercial coconut |
|-----|--|
| 573 | oil sample, the coconut oil possessed higher caprylic (9.4%), capric (6.3%), and medium |
| 574 | chain (69.7%) fatty acids. These fatty acid types are known to impart health benefits, and |
| 575 | contribute to higher oxidative stability to the oil itself. In addition, the resulting coconut oil |
| 576 | was also lower in acid value (0.27%) which also corresponds to lower free fatty acids, as |
| 577 | compared to the commercial coconut oil (0.91%). The free fatty acids are responsible for |
| 578 | undesirable flavour in the oil. Therefore overall, the coconut oil obtained from AEED |
| 579 | followed by chilling and thawing seems to possess greater oxidative stability, and the |
| 580 | attributes measured were within the Asian and Pacific Coconut Community standards |
| 581 | (Raghavendra & Raghavarao, 2010). |
| | |

582 Overall, enzyme based extraction methods result in oils with better characteristics 583 as compared to oil obtained from solvent and aqueous extraction methods. Therefore, 584 further studies are desirable to enable industrial application by scaling up.

585

586 5. Potentials for re-using enzymes in enzymatic extraction methods

587Rosenthal *et al.* (1996) highlighted the possible alternatives for improvement of aqueous

extraction, including the use of enzymes (i.e. AEE), the optimization of both extraction and

de-emulsification processes, utilization of membrane technology, and the potential of water

recycling (i.e. enzyme recycling in the case of AEE). Enzyme recycling may assist in

- reducing the cost of AEE which bears the potential to compete with conventional
- 592 extraction method based on the market price commanded by the oil (Nyam *et al.*, 2009a)

593 According to Jung et al. (2009), after conducting AEE (Protex 6L) to produce 594 soybean oil, the aqueous phase recovered contained 84.7% of the remaining Protex 6L 595 activity. After separation, a major part of this enzyme activity was recovered in the skim 596 fraction (Jung et al., 2009). Similarly, 100% of Protex 6L activity remained in the skim 597 fraction in a study conducted by Chabrand and Glatz (2009). These findings indicate the 598 possibility of recovering and re-using the skim fraction as a source of water and enzyme at the upstream end of the process (Jung et al., 2009). In addition, Jung et al. (2009) reported 599 600 lower Protex 6L activity in the cream emulsion, yet adequate to increase the free oil yield 601 with the use of suitable incubation time and temperature. Droplet coalescence was also 602 promoted by the gentle stirring during the incubation of the cream emulsion (Jung et al., 603 2009).

Studies concerning the enzyme recycling were conducted in order to improve 604 605 process economics and lower the environmental impact of the process. Another method 606 which has gained recent interests is the enzyme immobilization, where the enzymes are 607 separated from the treated products before being re-used. It was reported that the separated 608 enzymes possessed enhanced stability (Long et al., 2011; Wan et al., 2008; Roy et al., 2004). The increasing demands on enzyme-based methods have resulted in production of 609 610 more enzymes at lower production costs (Roy et al., 2004; Mondal et al., 2003; Sharma et 611 al., 2003; Chase, 1994).

612

613 6. Concluding remarks

614 This review has highlighted the main process, advantages, and disadvantages of AEE and 615 AEED as alternative methods for conventional solvent based extraction methods. In order 616 to enhance the oil yield, a combination of AEE with other non-enzymatic processing 617 methods prior to, or after AEE, has been widely conducted and relevant studies have been 618 reviewed in this paper. The process factors influencing AEE and AEED efficiencies, as 619 well as the oil characteristics, have also been discussed. On the whole, the process factors 620 are correlated with each other, and statistical optimization is currently the best solution for investigating the interacting effects between the contributing factors for obtaining highest 621 622 oil yield with favourable quality. The high cost of enzymes and production of lower oil 623 yield than that of solvent extraction method have been the major drawbacks of AEE process. Despite the problems, the interest in this method for oil and protein extraction has 624 625 progressively increased due to the perceived environmental advantages.

626

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Figure caption

| 921 | Fig. 1. Flow sheet for (a) production of extruded soybean oil by aqueous enzymatic |
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| 922 | extraction and free soybean oil recovery by aqueous enzymatic emulsion de-emulsification |
| 923 | method (Adapted from Lamsal and Johnson, 2007; Jung et al., 2009; Wu et al., 2009; |
| 924 | Chabrand and Glatz, 2009); (b) production of olive oil by aqueous enzymatic extraction |
| 925 | with different post-treatments (Adapted from Ranalli et al., 1999; Garcia et al., 2001; |
| 926 | Ranalli et al., 2001; Ranalli et al., 2003; De Faveri et al., 2008; Najafian et al., 2009); and |
| 927 | (c) production of virgin coconut oil by aqueous enzymatic emulsion de-emulsification |
| 928 | method (Adapted from Raghavendra and Raghavarao, 2010). |
| 020 | |

Table captions

Table 1. Commercial enzymes used for aqueous enzymatic extraction (AEE) and
aqueous enzymatic emulsion de-emulsification (AEED) processes: descriptions and
compositions.

935 Table 2. The oil yield enhancement with the use of enzymes, and the oil yield

936 difference between the enzyme and solvent extraction methods.

937 Table 3. Enhancement in oil yield due to presence of enzyme pre-treatment prior to

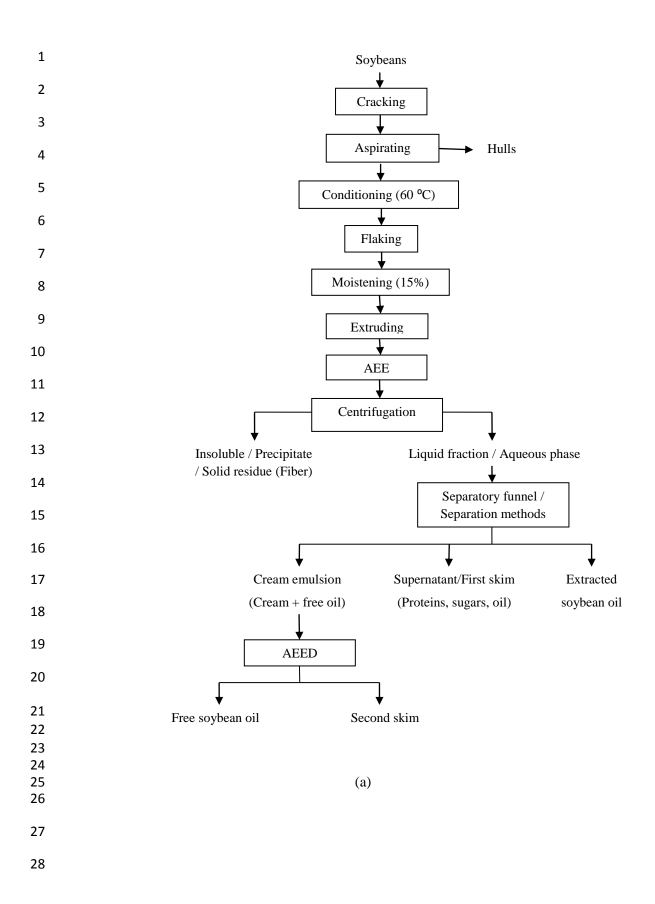
938 the extraction method, as compared to the extraction method alone.

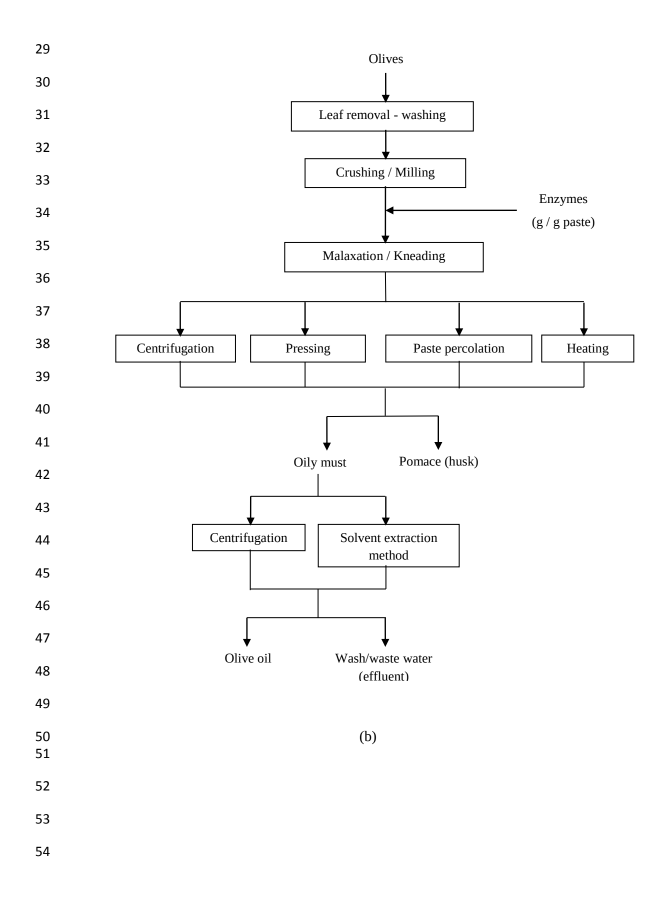
- 939 Table 4. The advantages of the use of pre-treatments (non-enzymatic) prior to the
- 940 enzymatic extraction method.
- 941 Table 5. Maximum oil yields as affected by the selected and optimized incubating
- 942 conditions of the aqueous enzymatic extraction and aqueous enzymatic emulsion de-
- 943 emulsification methods.
- 944 Table 6. The characteristics of oil yields from solvent, aqueous, and aqueous
- 945 enzymatic extraction methods.

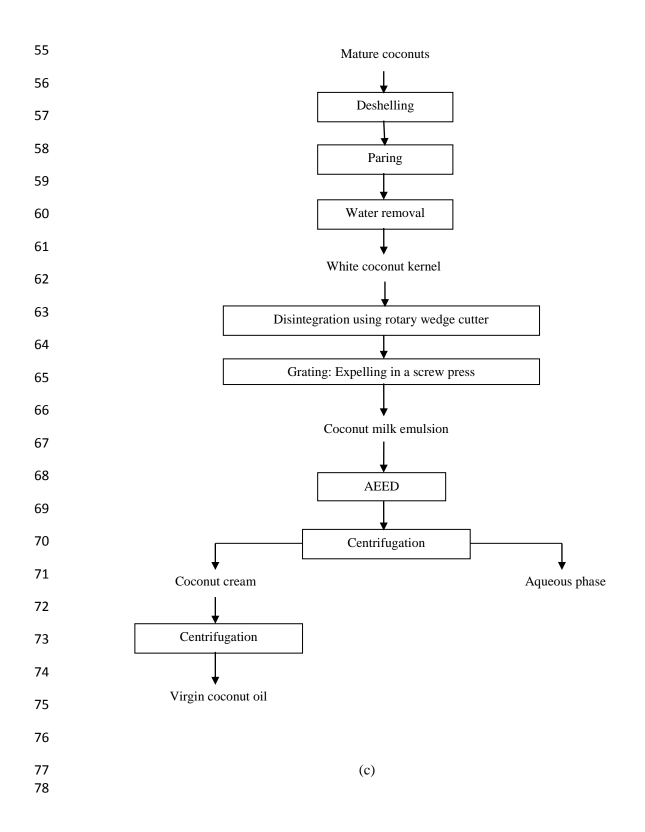
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| Enzymes commercial names | Description/Composition | Reference |
|---------------------------|----------------------------|----------------------------|
| | | |
| Single enzyme | | |
| Alcalase® | Protease | Womeni et al. (2008) |
| Alcalase 2.4L | Protease | Rosenthal et al. (2001) |
| | | Latif & Anwar (2009) |
| | | Jiang <i>et al.</i> (2010) |
| | | Latif & Anwar (2011) |
| | | Rovaris et al. (2012) |
| | | Tabtabaei & Diosady (2013) |
| As1398 | Protease | Jiang <i>et al.</i> (2010) |
| Celluclast 1.5L® | Cellulase | Dominguez et al. (1996) |
| | | Sineiro et al. (1998) |
| | | Abdulkarim et al. (2006) |
| | | Rovaris et al. (2012) |
| | | Tabtabaei & Diosady (2013) |
| | | Teixeira et al. (2013) |
| Flavourzyme® 1000 L | Protease | Nyam et al. (2009) |
| | | Nyam et al. (2009a) |
| Glucanex | Glucosidases | Garcia et al. (2001) |
| G-ZYME® G999 | Lysophospholipase A1 | Chabrand & Glatz (2009) |
| | | Wu et al. (2009) |
| | | Tabtabaei & Diosady (2013) |
| Lipomod 699L | Phospholipase A2 | Tabtabaei & Diosady (2013) |
| LysoMax TM | Phospholipase A2 | Wu et al. (2009) |
| Multifect Neutral® | Protease | Lamsal & Johnson (2007) |
| Neutrase 0.8L | Bacterial neutral protease | Abdulkarim et al. (2005) |
| | | Abdulkarim et al. (2006) |
| | | Nyam et al. (2009) |
| | | Nyam et al. (2009a) |
| Nutrase | Xylanase | Jiang et al. (2010) |
| Papain | Protease | Jiang et al. (2010) |
| Pectinase 1.06021 | Pectinase | Najafian et al. (2009) |
| Pectinase Multieffect FE® | Pectinase | Teixeira et al. (2013) |

 Table 1. Commercial enzymes used for aqueous enzymatic extraction (AEE) and aqueous enzymatic emulsion

 de-emulsification (AEED) processes: descriptions and compositions.

| Pectinex® | Pectinase | Womeni et al. (2008) |
|---------------------|---|----------------------------|
| Pectinex Ultra SP | Pectinase | Dominguez et al. (1996) |
| Pectinex Ultra SP-L | Pectinase | Abdulkarim et al. (2006) |
| | | Tabtabaei & Diosady (2013) |
| Promozyme | Pullulanase | Shah <i>et al.</i> (2005) |
| Protamex | Protease | Jiang et al. (2010) |
| Protex 6L | Alkaline serine endopeptidase | Chabrand & Glatz (2009) |
| | | Jung et al. (2009) |
| | | Wu et al. (2009) |
| | | Shan Liu et al. (2011) |
| | | Xiaonan Sui et al. (2011) |
| | | Tabtabaei & Diosady (2013) |
| Protex 7L | Natural metallo endopeptidase | Latif et al. (2008) |
| | | Chabrand & Glatz (2009) |
| | | Jung & Mahfuz (2009) |
| | | Latif & Anwar (2009) |
| | | Wu et al. (2009) |
| | | Latif & Anwar (2011) |
| | | Latif <i>et al.</i> (2011) |
| Protex 30L | Alkaline serine endopeptidase | Chabrand & Glatz (2009) |
| Protex 50FP | Acid fungal endopeptidase- | Wu et al. (2009) |
| | exopeptidase complex | |
| Protex 51FP | Neutral fungal endopeptidase- | Wu et al. (2009) |
| | exopeptidase complex | Tabtabaei & Diosady (2013) |
| Protex 89L | Endopeptidase | Tabtabaei & Diosady (2013) |
| ROHALASE® OS | Cellulase | Szydłowska-Czerniak et al. |
| ROHAPECT® PTE | Pectinase | (2010) |
| Termamyl 120L | α-amylase | Abdulkarim et al. (2006) |
| Enzymes mixture | | |
| Bioliva | Cellulase, hemicellulase, pectinase, | Ranalli et al. (2003) |
| | other minor enzymes | |
| Cytolase 0 | Cellulase, hemicellulase, pectinase, | Ranalli et al. (1999) |
| | other minor enzymes | Ranalli et al. (2003) |
| Kemzyme | Cellulase complex, hemi-cellulase | Latif & Anwar (2009) |
| | complex, α -amylase, β -glucanase, | Latif & Anwar (2011) |
| | protease, xylanase | Latif <i>et al.</i> (2011) |

| Maxoliva | Cellulase, hemicellulase, pectinase, | Ranalli et al. (2003) |
|-------------------------|---|-----------------------------|
| | other minor enzymes | |
| Multifect CX 13L | Cellulase, hemicellulase, β-glucanase, | Latif et al. (2008) |
| | arabinoxylans | Latif <i>et al.</i> (2011) |
| Multifect Pectinase FE | Pectinase, cellulase, hemicellulase | Latif et al. (2008) |
| Natuzyme | Cellulase, xylanase, phytase, α- | Latif et al. (2008) |
| | amylase, pectinase | Latif & Anwar (2009) |
| | | Latif & Anwar (2011) |
| | | Latif <i>et al.</i> (2011) |
| Olivex | Cellulase, hemicellulase, pectinase | Garcia et al. (2001) |
| Olivex-Celluclast | 50%: Cellulase, hemicellulase pectinase | Soto <i>et al.</i> (2007) |
| | 50%: Cellulase, hemicellulase | |
| Pectinex Ultra SP-L | Cellulase, pectinase, xylanase | Shah <i>et al.</i> (2005) |
| | | Najafian et al. (2009) |
| | | Tabtabaei & Diosady (2013 |
| Protizyme TM | Three different proteases with pH | Sharma <i>et al.</i> (2002) |
| | optima 3-4, 5-7, 7-10 | Gaur et al. (2007) |
| | | Jiang et al. (2010) |
| Rapidase® Liq plus | Hemicellulases, pectinases, cellulases | Gros et al. (2003) |
| Viscozyme® | (Carbohydrases): Cellulase, | Sowbhagya et al. (2009) |
| | hemicellulase, arabinase, xylanase, amylase, β-glucanase | Womeni et al. (2008) |
| Viscozyme L | (Carbohydrases): Cellulase, | Latif & Anwar (2009) |
| | hemicellulase, arabinase, xylanase, β- | Latif & Anwar (2011) |
| | glucanase | Latif <i>et al.</i> (2011) |
| | | Rovaris et al. (2012) |
| | | Tabtabaei & Diosady (2013 |

| Oil-bearing material | Type of enzyme | Difference in oil yield | Reference | |
|---|---------------------------------------|-------------------------|----------------------|-----------------------------|
| | | Aqueous extraction | Solvent treatment | _ |
| | | and aqueous | and aqueous | |
| | | enzymatic extraction | enzymatic extraction | |
| Crushed borage seeds (≤2.0 mm) | Olivex / Celluclast (1:1) | 7.80 | - | Soto et al. (2007) |
| Extruded soybean flakes | Protease | 20.00 | - | Lamsal <i>et al.</i> (2006) |
| | Multifect Neutral® | 13.40 | - | Lamsal & Johnson (2007) |
| | Protex 7L | 22.10 | - | Jung & Mahfuz (2009) |
| | Protex 51FP | 16.00 ^a | - | Wu et al. (2009) |
| | Protex 6L | 20.00 ^a | - | |
| | Protex 7L | 17.00 ^a | - | |
| Ground canola seeds | Multifect CX 13L | 9.50 | 17.10 | Latif et al. (2008) |
| | Protex 7L | 6.90 | 19.70 | |
| | Natuzyme | 6.20 | 20.40 | |
| Ground Jatropha seed kernels (inedible) | Protizyme TM | 26.00 | | Shah <i>et al.</i> (2005) |
| Ground Kalahari melon seeds | Neutrase 0.8L | | 9.58 | Nyam et al. (2009a) |
| | Flavourzyme 1000L | | 8.67 | |
| Ground Moringa. oleifera seeds | Neutrase 0.8L | | 8.20 | Abdulkarim et al. (2005) |
| | Neutrase 0.8L | 12.12 | 9.39 | Abdulkarim et al. (2006) |
| | Termamyl 120L | 10.15 | 11.36 | |
| | Pectinex Ultra SP-L | 6.98 | 14.53 | |
| | Celluclast 1.5L | 10.12 | 11.39 | |
| | Neutrase 0.8L / Termamyl 120L / | 12.83 | 8.68 | |
| | Pectinex Ultra SP-L / Celluclast 1.5L | | | |

| Table 2. Oil yield difference | e between the aqueous and aqueou | s enzymatic extraction, and between solvent and aq | ueous enzymatic extraction methods. |
|-------------------------------|----------------------------------|--|-------------------------------------|
| Oil-bearing material | Type of enzyme | Difference in oil vield (%) | Reference |

| | Natuzyme | 9.10 | 23.30 | Latif <i>et al.</i> (2011) |
|--------------------------------|-------------------------|-------|-----------|-----------------------------|
| | Kemzyme | 10.30 | 22.10 | |
| | Multifect CX 13L | 14.00 | 18.40 | |
| | Protex 7L | 14.70 | 17.70 | |
| | Viscozyme L | 13.10 | 19.30 | |
| Ground peanuts | Alcalase | 42.86 | - | Jiang et al. (2010) |
| | As1398 | 35.77 | - | |
| | Nutrase | 29.49 | - | |
| | Protizyme | 24.43 | - | |
| | Protamex | 18.30 | - | |
| | Protizyme TM | - | 3.36-5.88 | Sharma <i>et al.</i> (2002) |
| | Papain | - | 10.08 | |
| | Chymotrypsin | - | 16.38 | |
| | Trypsin | - | 13.86 | |
| Ground sesame seeds | Alcalase 2.4L | 12.50 | 25.40 | Latif & Anwar (2011) |
| | Natuzyme | 4.50 | 33.40 | |
| | Protex 7L | 6.40 | 31.50 | |
| | Viscozyme L | 9.10 | 28.80 | |
| | Kemzyme | 4.20 | 33.70 | |
| Ground sunflower seeds (0.75-1 | Celluclast 1.5L | 35.00 | - | Sineiro et al. (1998) |
| mm) | | | | |
| Ground sunflower seeds | Alcalase 2.4L | 8.30 | 18.90 | Latif & Anwar (2009) |
| | Kemzyme | 13.90 | 13.30 | |
| | Natuzyme | 17.20 | 10.00 | |
| | Protex 7L | 10.00 | 17.20 | |
| | Viscozyme L | 21.40 | 5.80 | |
| Heat-treated soybean flour | Alcalase 2.4L | 16.90 | - | Rosenthal et al. (2001) |

| Kernel flour of bush mango | Alcalase® | 7.60 | - | Womeni et al. (2008) |
|---------------------------------|--|----------------------------|-------|-------------------------------|
| | Pectinex® | 14.80 | - | |
| | Viscozyme® | 40.60 | - | |
| Minced yellow horn seed kernels | Cellulase / Hemicellulase / Pectinase | | 9.00 | Li et al. (2013) |
| | (1.8:1.3:2.5) | | | |
| Olive paste | Bioliva | 1.20 | - | Ranalli et al. (2003) |
| | Maxoliva | 1.37 | - | |
| | Cytolase 0 | 1.44 | - | |
| | A (pectinase, cellulase, hemicellulase) | 152.00 (30 min) | - | Aliakbarian et al. (2008) |
| | / B (pectinase, hemicellulase) / | ulase) / 91.40 (150 min) - | | |
| | C (pectolytic enzyme) (1:1:1) | 91.40 (150 mm) | - | |
| | Pectinex Ultra SP-L | 1.96 ^b | - | Najafian <i>et al.</i> (2009) |
| | Pectinase 1.6021 | 1.41 ^b | - | |
| Palm fruit | Pectinase / cellulase | 35.57 | 5.36 | Teixeira et al. (2013) |
| | Pectinase / cellulase / tannase | 35.90 | 5.03 | |
| | Tannase | 12.70 | 28.23 | |
| Rapeseed slurry | Pectinase | 38.10 | - | Zhang <i>et al.</i> (2007) |
| | Cellulase | 21.50 | - | |
| | B-glucanase | 16.20 | - | |
| | Pectinase / Cellulase / β -glucanase | 43.80 | - | |
| | (4:1:1) | | | |
| | Multifect Pectinae FE | 5.70 | - | |
| Shattered bayberry kernels (60- | Cellulase / Neutral protease (1:2) | | 31.85 | Zhang <i>et al.</i> (2012) |
| mesh sieved) | | | | |
| Yellow mustard flour | Celluclast 1.5L | 3.74 | 10.59 | Tabtabaei & Diosady |
| | Pectinex Ultra SP-L | 3.03 | 11.30 | (2013) |

| Viscozyme L | 3.99 | 10.34 |
|---|------|-------|
| Celluclast 1.5L / Pectinex Ultra SP-L / | 6.70 | 7.63 |
| Viscozyme L (1: 1:1) | | |

The oil yield differences were determined based on the oil yields under the best incubating conditions of each enzyme used, or based on the fixed incubating conditions for all enzymes used, in the conducted studies.

All aqueous enzymatic extractions resulted in higher oil yields than aqueous extractions, and all solvent treatments resulted in higher oil yields than aqueous enzymatic extractions.

^a total oil as in the skim and cream emulsion

^b average oil yield enhancements from three olive species with the use of enzymes at high concentrations

| Oil-bearing material | Type of enzyme | Extraction method | Enhancement | Reference |
|-------------------------|-----------------------------|------------------------------|-------------------|-----------------------|
| | (pre-treatment) | | in oil yield | |
| | | | (%) | |
| Crushed borage seeds | Olivex / Celluclast (1:1) | Double pressing | 5.40 ^a | Soto et al. |
| (≤2.0 mm) | | | | (2007) |
| Crushed garlic cloves | Cellulase | Steam distillation | 0.11 | Sowbhagya e |
| | Pectinase | | 0.23 | al. (2009) |
| | Protease | | 0.22 | |
| | Viscozyme | | 0.18 | |
| | Cellulase | Hydrodistillation | 0.14 | |
| | Pectinase | | 0.26 | |
| | Protease | | 0.24 | |
| | Viscozyme | | 0.19 | |
| Ground flaxseeds | Cellulase / Pectinase / | Ultrasonication | 29.50 | Long et al. |
| | Hemicellulase (1:1:1) | | | (2011) |
| Ground rapeseeds | ROHAPECT® PTE | Pressing | 5.70 | Szydłowska- |
| | | | 1 70 | Czerniak et |
| | ROHALASE® OS | | 1.70 | al. (2010) |
| Milled grape seeds | A mixture of cellulase, | Solvent extraction (24 hr) | 106.00 | Passos et al. |
| | xylanase, protease. | | 162.00 | (2009) |
| | pectinase | Solvent extraction (120 hr) | 163.00 | |
| Minced yellow horn seed | Cellulase / hemicellulase / | Microwave | 4.30 (oil yield | Li et al. |
| kernels | pectinase (1.8 : 1.3 : 2.5) | | enhancement | (2013) |
| | | | as compared to | |
| | | | AEE alone) | |
| Pre-heated ground | Ultrazyme / Celluclast | Double pressing (hydraulic | ~8.00 | Zuniga <i>et al</i> . |
| Chilean hazelnut seeds | (1:1) | pressing at each of 39.2 | | (2003) |
| (inedible, ≤1.4 mm) | | MPa) | | |
| Silybum marianum seed | Cellulase / Xylanase / | Solvent extraction (1.5 hr) | 10.46 | Li <i>et al</i> . |
| powders | Pectinase / Protease | | 50.72 | (2012) |
| | (2:1:1:2) | Solvent extraction (14.0 hr) | 50.72 | |
| Whole sunflower kernels | Celluclast 1.5L / Pectinex | Pressing (Batch press) | 13.11 | Dominguez e |
| | Ultra SP (2:1) | | | al. (1996) |
| Mango kernel powders | Protizyme TM | Three-phase partitioning | 16.00 | Gaur et al. |
| Soybean flour | | method | 8.00 | (2007) |

 Table 3. Enhancement in oil yield due to presence of enzyme pre-treatment prior to the extraction method, as

 compared to the extraction method alone.

| Rice bran powders | 14.00 |
|-------------------|-------|

^a the oil yield enhancement was based on the difference between an enzymatic and non-enzymatic pre-treatment,

followed by double pressing

| Oil-bearing | Pre-treatment | Type of | Advantages | Reference |
|---------------|-----------------|-------------------------|---|---------------------|
| material | | enzyme | | |
| Ground Isatis | Microwave | Cellulase / | - In combination with AEE, the use of | Gai et al. (2013) |
| indigotica | | Proteinase / | optimal microwave irradiation power | |
| seeds | | Pectinase | increased the oil yield up to 59.27%, and | |
| | | (1:1:1) | the oil yield had greater antioxidant | |
| | | | properties than solvent-extracted oil. | |
| Ground | Ultrasonication | Protizyme TM | The enzyme treatment time was reduced | Shah <i>et al</i> . |
| Jatropha seed | (5 min) | | from 18 hr to 6 hr for maximum of 74% oil | (2005) |
| kernels | | | yield | |
| (inedible) | | | | |
| Ground | Electrical | - | Mucilage (stabilizing agent) is removed | Gros et al. (2003 |
| linseeds | discharge | | which caused easier oil separation from the | |
| | | | resulted residue by using enzyme treatment | |
| Grounds | Alkaline | Alcalase | Oil yield of 5.87% higher than AEE alone | Jiang et al. |
| peanuts | extraction | | | (2010) |
| Ground pitaya | Microwave | Pectinase / | - Oil yield of 0.84% higher than AEE | Rui et al. (2009) |
| seeds (40- | | Cellulase / | alone | |
| mesh sieved) | | Acid protease | | |
| | | (1:1:1) | | |
| Ground | Ultrasound | Protex 6L | -Under the fixed parameters of the | Xiaonan Sui et |
| watermelon | | | ultrasound, the yield was 20.67% higher | al. (2011), Shan |
| kernels | | | than AEE alone | Liu et al. (2011) |
| | | | -Under the selected parameters of | |
| | | | ultrasound for maximum oil yield, the | |
| | | | yield was 21.39% higher than AEE alone | |
| Soybean | High pressure | Protex 7L | Oil yield of 3.20% higher than AEE alone | Jung & Mahfuz |
| flakes | processing (200 | | | (2009) |
| | MPa) | | | |
| | High pressure | | Oil yield of 1.30% higher than AEE alone | |
| | processing (500 | | | |
| | MPa) | | | |
| | Extrusion | | - Oil yield of 29.90% higher than AEE | |
| | | | alone | |
| | | | - Free oil yield of 17.00% higher than AEE | |

Table 4. The advantages of the use of pre-treatments (non-enzymatic) prior to the enzymatic extraction method.

| | | alone | |
|-----------|-----------|--|--------------------|
| Extrusion | Protex 6L | - Oil yield of 35.52% higher than AEE | Jung et al. (2009) |
| | | alone | |
| | | - After de-emulsification: Free oil from | |
| | | cream emulsion of 62.00% higher than | |
| | | AEE alone | |

AEE: aqueous enzymatic extraction.

Table 5. Maximum oil yields as affected by the selected and optimized incubating conditions of the aqueous enzymatic extraction and aqueous enzymatic emulsion de-emulsification methods.

| Oil-bearing material | Type of enzyme | Moisture / | Enzyme / | pН | Tempera- | Time (hr) | Agitation | Oil yield | Reference |
|----------------------|----------------|----------------|----------|----|-----------|-----------|------------|-----------|-----------|
| | | Material ratio | Material | | ture (°C) | | rate (rpm) | (%) | |
| | | (w/w; for | ratio | | | | | | |
| | | aqueous | | | | | | | |
| | | enzymatic | | | | | | | |
| | | extraction) | | | | | | | |

Selected(*) and optimized (**) incubating conditions used for maximum oil yield in aqueous enzymatic extraction

| Crushed borage seeds (≤2.0 mm) | Olivex / Celluclast (1:1) ^a | 20%* (corresponded to 1:5) | 0.25%* | - | 45.0* | 9.00* | - | 85.50 | Soto <i>et al.</i> (2007) |
|---|---|----------------------------------|-----------------|---------|-------|--------|------|-------|-------------------------------|
| Ground Jatropha seed kernels (inedible) | Protizyme ^{TM a} | 6:1 | 0.25 (w/w)% | 9.00* | 50.0* | 18.00 | 100 | 64.00 | Shah <i>et al.</i> (2005) |
| Ground Moringa. | Celluclast 1.5L ^a | 6:1 | 2.00% | 4.80*** | 60.0* | 36.00* | 120* | 22.01 | Abdulkarim et |
| oleifera seeds | Termamyl 120L ^a | | (v/w)* | 5.50*** | | | | 22.04 | al. (2006) |
| | Pectinex Ultra SP-L ^a | | | 3.50*** | 45.0* | | | 18.87 | |
| | Neutrase 0.8L ^a | | | 6.80*** | | | | 24.02 | |
| | Neutrase 0.8L / | | | 7.50*** | | | | 24.72 | |
| | Termamyl 120L / | | | | | | | | |
| | Pectinex Ultra SP-L / | | | | | | | | |
| | Celluclast 1.5L ^a | | | | | | | | |
| Ground peanuts | Alcalase ^a | 5:1* | 1.50% (w/w)* | 8.50* | 60.0* | 5.00* | - | 73.45 | Jiang <i>et al.</i> (2010) |

| | Protizyme ^{TM a} | 2:1 | 2.50% | 4.00* | 40.0* | 18.00* | 80* | 36.12- | Sharma et al. |
|----------------------|---|-----------|---------|----------|---------|---------|------------|--------|----------------------|
| | | | (w/w)* | | | | | 38.64 | (2002) |
| Ground pitaya seeds | Pectinase / Cellulase / | 8:1 | - | 7.00 | 50.0* | 1.00 | 90 | 6.94 | Rui et al. |
| (40-mesh sieved) | Acid protease (1:1:1) a | | | | | | | | (2009) |
| Ground rice bran | Alcalase 0.6L ^a | - | 1.00% | 9.00 | 60.0* | 3.00* | 1000 | 79.10 | Hanmoungjai |
| (16-mesh sieved) | | | (w/w)* | | | | | | et al. (2001) |
| Ground sunflower | Celluclast 1.5L ^a | 5:1* | 2.00% | 4.80*** | 50.0*** | 2.00* | 150 | 35.65 | Sineiro et al. |
| seeds (0.75-1 mm) | | | (w/w)* | | | | | | (1998) |
| Heat-treated soybean | Alcalase 2.4L ^a | - | 3.00% | 8.00 *** | 50.0*** | 1.00 | 200 | 58.70 | Rosenthal et |
| flour | | | (v/w)* | | | | | | al. (2001) |
| Olive paste | A (pectinase, | - | 0.25% | - | 30.0 | 2 hr 30 | 10 | 17.50 | Aliakbarian et |
| | cellulase, | | (v/w)* | | | min* | (kneading) | | al. (2008) |
| | hemicellulase) / | | | | | | | | |
| | B (pectinase, | | | | | | | | |
| | hemicellulase) / | | | | | | | | |
| | C (pectolytic | | | | | | | | |
| | enzyme) (1:1:1) ^a | | | | | | | | |
| Rapeseed slurry | Pectinase / Cellulase / | 5:1* | 2.50% | 5.00 | 48.0 | 4.00* | 200 | 92.70 | Zhang <i>et al</i> . |
| | β -glucanase (4:1:1) ^a | | (v/w)* | | | | | | (2007) |
| Ground Kalahari | Neutrase 0.8L ^a | - | 2.50% | 7.00** | 58.0** | 31.00** | 100 | 68.58 | Nyam et al. |
| melon seeds | | | (w/w)** | | | | | | (2009a) |
| | Flavourzyme® 1000 | - | 2.10% | 6.00** | 50.0** | 36.00** | 100 | 71.55 | |
| | La | | (w/w)** | | | | | | |
| Ground Moringa. | Neutrase 0.8L ^a | 6:1 (v/w) | 2.00% | 6.80 *** | 45.0** | 24.00** | 120 | 22.60 | Abdulkarim et |
| oleifera seeds | | | (v/w) | | | | | | al. (2005) |

| Ground pine kernels | Alcalase endo- | 5:1** | 1.97%** | 8.40** | 51.0** | 3.00** | - | 89.12 | Yang Li <i>et al.</i> |
|---------------------|------------------------------|----------------|---------|--------|--------|--------|-----|-------|-------------------------|
| a | protease ^a | | | | | | | 00.10 | (2011) |
| Ground pumpkin | Cellulase ^a | - | 1.70% | - | 47.0** | 2.64** | - | 89.12 | Hu & Zou |
| seeds | | | (w/w)** | | | | | | (2013) |
| Ground watermelon | Protex 6L ^a | 4.35:1** | 2.63%** | 7.89** | 47.1** | 4.29** | - | 77.25 | Xiaonan Sui et |
| kernels | | | | | | | | | al. (2011); |
| | | | | | | | | | Shan Liu <i>et al</i> . |
| | | | | | | | | | (2011) |
| Palm fruits | Pectinase / Cellulase / | 2:1 (v/w)** | 4.00** | 4.00** | 50.0 | 0.50* | 200 | 91.52 | Teixeira et al. |
| | Tannase (1:1:1) ^a | | | | | | | | (2013) |
| Shattered bayberry | Cellulase / Neutral | 4.91:1 (v/w)** | 3.17%** | - | 51.6** | 4.00** | - | 31.15 | Zhang et al. |
| kernels (60-mesh | protease (1:2) ^a | | | | | | | | (2012) |
| sieved) | | | | | | | | | |
| | | | | | | | | | |

Selected (*) and optimized (**) incubating conditions for maximum free oil yield in aqueous enzymatic emulsion de-emulsification method

| Alkaline pre-treated | Alcalase 2.4L ^a | As1398 ^b | 1.00% | - | - | 2.0 hr | - | 12.66 | Jiang <i>et al</i> . |
|----------------------|----------------------------|-----------------------------|-------|--------|---------|---------|---|-------|----------------------|
| ground peanuts | | | | | | | | | (2010) |
| Coconut milk | - | Aspartic | 0.10% | - | 37.0* | 3.0 hr | - | 83.00 | Raghavendra |
| emulsion | | protease | | | | | | | & Raghavarao |
| | | (endoprotease) ^b | | | | | | | (2010) |
| Extruded soybean | Protease Multifect | LysoMax TM / | - | 4.5*** | 60.0*** | 1 hr 30 | - | 68.00 | Lamsal & |
| flakes | Neutral® ^a | G-ZYME G999 | | | | min | | | Johnson (2007) |
| | | (1:1) ^b | | | | | | | |
| | | Phospholipase | - | 7.0*** | 37.0*** | 1 hr 30 | - | 73.00 | |
| | | | | | | | | | |

| | | C ^b | | | | min | | | |
|----------------------|------------------------|-------------------------------|------------------|----------------|--------------|----------|------|--------|----------------------------|
| | Protex 6L ^a | Protex 6L ^b | 2.50%* | 4.5* | 50.0 | 1 hr 30 | - | 100.00 | de Moura et al |
| | | | | | | min | | | (2008) |
| | Protex 6L ^a | Protex 6L ^b | 1.25%** | - | 50.0** | 1 hr 30 | - | 100.00 | Jung et al. |
| | | | | | | min** | | | (2009) |
| | Protex 7L ^a | LysoMax ^{TM b} | 2.00% | 8.0*** | 40.0*** | 1 hr 30 | - | 100.00 | Wu et al. |
| | | G-ZYME® G999 ^b | | 4.5*** | 50.0*** | min | | | (2009) |
| | | Protex 6L ^b | | 8.0*** | 50.0*** | | | | |
| | | Protex 7L ^b | | 7.0*** | 50.0*** | | | | |
| | | Protex 50FP ^b | | 4.5*** | 50.0*** | | | | |
| | | Protex 51FP ^b | | 8.0*** | 50.0*** | | | | |
| Ground Perilla | - | Protex 6L ^b | 1.90%** | 9.4** | 62.6** | 1.6 hr** | - | 85.52 | Zhang et al. |
| frutescens seeds | | | 0 000 / 1 | 4 ministration | 7 0 0 | 2 0 1 | | 100.00 | (2013) |
| Soybean flour | Protex 7L ^a | G-ZYME G999 ^b | 2.00%* | 4.5*** | 50.0 | 3.0 hr | 700* | 100.00 | Chabrand & Glatz (2009) |
| | | Protex 6L ^b | 3.00%* | 9.0*** | 50.0 | 3.0 hr | 500* | 72.00 | |
| Yellow mustard flour | Celluclast 1.5L/ | Protex 6L ^b | 2.50% | 4.5- | 50-60*** | 3.0 hr | - | 91.30 | Tabtabaei & |
| | Viscozyme L / | | | 6.0*** | | | | | Diosady |
| | Pectinex Ultra SP-L | Alcalase 2.4L ^b | | 6.5- | 45-65*** | | | 42.10 | (2013) |
| | (1:1:1) ^a | | | 8.5*** | | | | | |
| | | Lipomode 699L ^b | | 8.0*** | 40.0*** | | | 1.30 | |

| G-ZYME | 4.5*** | 50-60*** | 41.20 |
|-------------------|--------|----------|-------|
| G999 ^b | | | |

Values without any notation are fixed incubating conditions.

^a Type of enzymes used for aqueous enzymatic extraction

^b Type of enzymes used for aqueous enzymatic emulsion de-emulsification

*selected incubating condition; the authors varied the level of each incubating condition and finalized the conditions which resulted in highest oil yield.

**optimized incubating condition; the authors varied the level of each incubating condition and optimized the conditions which resulted in highest oil yield based on an experimental design and statistical software used.

*** optimum incubating condition of the enzyme used; different types of enzymes possess different optimum pH and temperature where the enzymes attain maximum activity

| Oil | Oil-bearing material | Solvent | Aqueous | Aqueou | is enzymatic extraction | Reference |
|--------------|-----------------------|------------|------------|--------|-------------------------|----------------------|
| characteris- | | extraction | extraction | | | |
| tic | | | | | | |
| Free fatty | Extruded soybean | 0.26 | * | 0.18 | Protex 6L | Jung <i>et al</i> . |
| acids (%) | flakes | 0.20 | | 0.10 | | (2009) |
| | Ground canola seeds | 0.81 | 0.56 | 0.52 | Multifect CX 13L | Latif <i>et al</i> . |
| | | | | 0.57 | Protex 7L | (2008) |
| | | | | 0.55 | Natuzyme | |
| | | | | 0.54 | Multifect Pectinae FE | |
| | Ground Kalahari | 0.60 | * | 0.90 | Flavourzyme® 1000 L | Nyam <i>et al</i> . |
| | melon seeds | | | 0.90 | Neutrase 0.8L | (2009) |
| | Ground Moringa. | 2.48 | * | 1.13 | Neutrase 0.8L | Abdulkarim |
| | oleifera seeds | | | | | et al. (2005) |
| | | 2.48 | 1.22 | 1.13 | Neutrase 0.8L | Abdulkarim |
| | | | | 1.24 | Termamyl 120L | et al. (2006) |
| | | | | 1.22 | Pectinex Ultra SP-L | |
| | | | | 1.25 | Celluclast 1.5L | |
| | | | | 1.23 | Neutrase 0.8L / | |
| | | | | | Termamyl 120L / | |
| | | | | | Pectinex Ultra SP-L / | |
| | | | | | Celluclast 1.5L | |
| | | 1.26 | 0.42 | 0.43 | Natuzyme | Latif et al. |
| | | | | 0.41 | Kemzyme | (2011) |
| | | | | 0.39 | Multifect CX 13L | |
| | | | | 0.38 | Protex 7L | |
| | | | | 0.42 | Viscozyme L | |
| | Ground rice bran (16- | 7.40 | * | 2.36 | Alcalase 0.6L | Hanmoungja |
| | mesh sieved) | | | | | et al. (2001) |
| | Ground sesame seeds | 0.54c | 0.48 | 0.47 | Natuzyme | Latif & |
| | | | | 0.44 | Kemzyme | Anwar (2011 |
| | | | | 0.51 | Protex 7L | |
| | | | | 0.46 | Alcalase 2.4L | |
| | | | | 0.44 | Viscozyme L | |
| | Ground sunflower | 0.94 | 0.68 | 0.66 | Alcalase 2.4L | Latif & |
| | seeds | | | 0.65 | Kemzyme | Anwar (2009 |

Table 6. The characteristics of oil yields from solvent, aqueous, and aqueous enzymatic extraction methods.

| | | | | 0.67 | Natuzyme | |
|--------------|-----------------------|--------|--------|--------|-------------------------|----------------------|
| | | | | 0.69 | Protex 7L | |
| | | | | 0.64 | Viscozyme L | |
| Iodine value | Ground canola seeds | 117.00 | 114.00 | 116.00 | Multifect CX 13L | Latif <i>et al</i> . |
| (g / 100g) | | | | 114.00 | Protex 7L | (2008) |
| | | | | 117.00 | Natuzyme | |
| | | | | 116.00 | Multifect Pectinae FE | |
| | Ground flaxseeds | 140.80 | * | 161.20 | Cellulase / Pectinase / | Long et al. |
| | | | | | Hemicellulase (1:1:1) | (2011) |
| | Ground Kalahari | 125.00 | * | 141.00 | Flavourzyme® 1000 L | Nyam et al. |
| | melon seeds | | | 135.20 | Neutrase 0.8L | (2009) |
| | Ground Moringa. | 65.40 | * | 66.10 | Neutrase 0.8L | Abdulkarim |
| | oleifera seeds | | | | | et al. (2005) |
| | | 65.40 | 66.00 | 67.10 | Neutrase 0.8L | Abdulkarim |
| | | | | 66.50 | Termamyl 120L | et al. (2006) |
| | | | | 67.20 | Pectinex Ultra SP-L | |
| | | | | 66.50 | Celluclast 1.5L | |
| | | | | 67.00 | Neutrase 0.8L / | |
| | | | | | Termamyl 120L / | |
| | | | | | Pectinex Ultra SP-L / | |
| | | | | | Celluclast 1.5L | |
| | | 67.00 | 70.00 | 76.00 | Natuzyme | Latif <i>et al</i> . |
| | | | | 73.00 | Kemzyme | (2011) |
| | | | | 75.00 | Multifect CX 13L | |
| | | | | 74.00 | Protex 7L | |
| | | | | 76.00 | Viscozyme L | |
| | Ground pitaya seeds | 173.10 | * | 118.00 | Pectinase / Cellulase / | Rui et al. |
| | (40-mesh sieved) | | | | Acid protease (1:1:1) | (2009) |
| | Ground rice bran (16- | 95.40 | * | 97.18 | Alcalase 0.6L | Hanmoungja |
| | mesh sieved) | | | | | <i>et al.</i> (2001) |
| | Ground sesame seeds | 107.00 | 106.00 | 104.00 | Natuzyme | Latif & |
| | | | | 109.00 | Kemzyme | Anwar (2011 |
| | | | | 108.00 | Protex 7L | |
| | | | | 105.00 | Alcalase 2.4L | |
| | | | | 103.00 | Viscozyme L | |
| | Ground sunflower | 127.00 | 120.00 | 124.00 | Alcalase 2.4L | Latif & |

| | seeds | | | 121.00 | Kemzyme | Anwar (2009) |
|----------------------|-----------------------|--------|------|--------|-------------------------|----------------------|
| | | | | 123.00 | Natuzyme | |
| | | | | 122.00 | Protex 7L | |
| | | | | 121.00 | Viscozyme L | |
| Peroxide | Extruded soybean | 6.50 | * | 4.05 | Protex 6L | Jung et al. |
| value (meq | flakes | | | | | (2009) |
| O ₂ / kg) | Ground canola seeds | 1.29 | 0.69 | 0.72 | Multifect CX 13L | Latif <i>et al</i> . |
| | | | | 0.70 | Protex 7L | (2008) |
| | | | | 0.71 | Natuzyme | |
| | | | | 0.64 | Multifect Pectinae FE | |
| | Ground flaxseeds | 1.20 | * | 1.00 | Cellulase / Pectinase / | Long et al. |
| | | | | | Hemicellulase (1:1:1) | (2011) |
| | Ground Kalahari | 2.30 | * | 6.40 | Flavourzyme® 1000 L | Nyam <i>et al</i> . |
| | melon seeds | | | 7.30 | Neutrase 0.8L | (2009) |
| | Ground Moringa. | 2.09 | 1.60 | 1.58 | Natuzyme | Latif <i>et al</i> . |
| | oleifera seeds | | | 1.56 | Kemzyme | (2011) |
| | | | | 1.61 | Multifect CX 13L | |
| | | | | 1.63 | Protex 7L | |
| | | | | 1.59 | Viscozyme L | |
| | Ground pitaya seeds | 1.93 | * | 1.44 | Pectinase / Cellulase / | Rui et al. |
| | (40-mesh sieved) | | | | Acid protease (1:1:1) | (2005) |
| | Ground rice bran (16- | 8.20 | * | 12.01 | Alcalase 0.6L | Hanmoungjai |
| | mesh sieved) | | | | | et al. (2001) |
| | Ground sesame seeds | 1.50 | 1.30 | 0.90 | Natuzyme | Latif & |
| | | | | 1.30 | Kemzyme | Anwar (2011) |
| | | | | 1.40 | Protex 7L | |
| | | | | 1.10 | Alcalase 2.4L | |
| | | | | 1.20 | Viscozyme L | |
| | Ground sunflower | 1.78 | 1.36 | 1.25 | Alcalase 2.4L | Latif & |
| | seeds | | | 1.33 | Kemzyme | Anwar (2009) |
| | | | | 1.32 | Natuzyme | |
| | | | | 1.31 | Protex 7L | |
| | | | | 1.37 | Viscozyme L | |
| Saponifica- | Ground Kalahari | 173.20 | * | 185.20 | Flavourzyme® 1000 L | Nyam <i>et al</i> . |
| tion value | melon seeds | | | 184.80 | Neutrase 0.8L | (2009) |

| (mg KOH / g | Ground Moringa. | 164.00 | * | 163.00 | Neutrase 0.8L | Abdulkarim |
|---------------------------------------|-----------------------|--------|--------|--------|-------------------------|----------------------|
| oil) | oleifera seeds | | | | | et al. (2005) |
| | | 164.00 | 158.00 | 156.00 | Natuzyme | Latif <i>et al</i> . |
| | | | | 158.00 | Kemzyme | (2011) |
| | | | | 155.00 | Multifect CX 13L | |
| | | | | 159.00 | Protex 7L | |
| | | | | 156.00 | Viscozyme L | |
| | Ground pitaya seeds | 194.40 | * | 191.10 | Pectinase / Cellulase / | Rui et al. |
| | (40-mesh sieved) | | | | Acid protease (1:1:1) | (2005) |
| | Ground rice bran (16- | 187.60 | * | 188.72 | Alcalase 0.6L | Hanmoungjai |
| | mesh sieved) | | | | | et al. (2001) |
| | Ground sesame seeds | 169.00 | 159.00 | 158.00 | Natuzyme | Latif & |
| | | | | 162.00 | Kemzyme | Anwar (2011) |
| | | | | 167.00 | Protex 7L | |
| | | | | 164.00 | Alcalase 2.4L | |
| | | | | 156.00 | Viscozyme L | |
| | Ground sunflower | 190.00 | 187.00 | 187.00 | Alcalase 2.4L | Latif & |
| | seeds | | | 186.00 | Kemzyme | Anwar (2009) |
| | | | | 187.00 | Natuzyme | |
| | | | | 187.00 | Protex 7L | |
| | | | | 185.00 | Viscozyme L | |
| Total | Ground canola seeds | 739.00 | 598.00 | 794.00 | Multifect CX 13L | Latif <i>et al</i> . |
| tocopherols; | | | | 805.00 | Protex 7L | (2008) |
| α , δ , and γ | | | | 783.00 | Natuzyme | |
| $(\alpha, \beta, \delta, and \gamma)$ | | | | 819.00 | Multifect Pectinae FE | |
| for Kalahari | Ground Kalahari | 174.80 | * | 143.20 | Flavourzyme® 1000 L | Nyam <i>et al</i> . |
| melon seeds | melon seeds | | | 143.30 | Neutrase 0.8L | (2009) |
| and olive | Ground Moringa | 179.30 | 216.90 | 220.80 | Natuzyme | Latif <i>et al</i> . |
| paste) | oleifera seeds | | | 228.50 | Kemzyme | (2011) |
| (mg / kg oil) | | | | 221.70 | Multifect CX 13L | |
| | | | | 221.50 | Protex 7L | |
| | | | | 228.30 | Viscozyme L | |
| | Ground sesame seeds | 584.10 | 603.30 | 628.50 | Natuzyme | Latif & |
| | | | | 641.20 | Kemzyme | Anwar (2011) |
| | | | | 627.30 | Protex 7L | |
| | | | | 619.80 | Alcalase 2.4L | |

| | _ | _ | | | 612.80 | Viscozyme L | |
|----------------|------------------|--------------|--------|--------|--------|-------------------------|------------------------|
| | Ground sunflower | | 799.00 | 778.00 | 845.00 | Alcalase 2.4L | Latif & |
| | seeds | | | | 849.00 | Kemzyme | Anwar (2009 |
| | | | | | 849.00 | Natuzyme | |
| | | | | | 842.00 | Protex 7L | |
| | | | | | 833.00 | Viscozyme L | |
| | Olive | Cipressino | * | 77.30 | 89.20 | Cytolase 0 | Ranalli <i>et al</i> . |
| | paste | Cassanese | | 95.20 | 114.10 | | (2001) |
| | | Leccino | | 117.00 | 135.40 | | |
| | | Dritta | * | 231.00 | 288.00 | Cytolase 0 | Ranalli <i>et al</i> . |
| | | | | | 279.00 | Maxoliva | (2003) |
| | | | | | 266.00 | Bioliva | |
| | | Caroleo | * | 218.00 | 273.00 | Cytolase 0 | |
| | | | | | 269.00 | Maxoliva | |
| | | | | | 252.00 | Bioliva | |
| | | Coratina | * | 244.00 | 305.00 | Cytolase 0 | |
| | | | | | 300.00 | Maxoliva | |
| | | | | | 289.00 | Bioliva | |
| | Palm fru | uit | * | 325.27 | 251.11 | Pectinase / Cellulase | Teixeira et al |
| | | | | | 200.54 | Pectinase / Cellulase / | (2013) |
| | | | | | | Tannase | |
| | | | | | 204.26 | Tannase | |
| Total | Ground | Kalahari | 18.00 | * | 18.00 | Flavourzyme® 1000 L | Nyam <i>et al</i> . |
| phenolic | melon s | eeds | | | 19.00 | Neutrase 0.8L | (2009) |
| content | Ground | Moringa | 12.00 | 13.00 | 15.00 | Natuzyme | Latif <i>et al</i> . |
| mg / kg oil), | oleifera | seeds | | | 14.00 | Kemzyme | (2011) |
| as in gallic | | | | | 13.00 | Multifect CX 13L | |
| acid | | | | | 14.00 | Protex 7L | |
| equivalent for | | | | | 18.00 | Viscozyme L | |
| sesame seeds, | Ground | sesame seeds | 17.00 | 18.00 | 19.00 | Natuzyme | Latif & |
| sunflower | | | | | 18.00 | Kemzyme | Anwar (2011 |
| seeds, | | | | | 22.00 | Protex 7L | |
| Moringa | | | | | 21.00 | Alcalase 2.4L | |
| oleifera | | | | | 24.00 | Viscozyme L | |
| seeds, and | Ground | sunflower | 8.00 | 9.00 | 13.00 | Alcalase 2.4L | Latif & |
| palm fruit; | seeds | | | | 14.00 | Kemzyme | Anwar (2009 |

| caffeic acid | | | | | 13.00 | Natuzyme | |
|----------------|-------|------------|-------|--------|-----------------------|-------------------------|------------------------|
| equivalent for | | | | | 13.00 | Protex 7L | |
| olive paste; | | | | | 15.00 | Viscozyme L | |
| and sum of | Olive | Cipressino | * | 90.00 | 105.00 | Cytolase 0 | Ranalli et al. |
| phenolic | paste | Cassanese | | 122.00 | 153.00 | | (2001) |
| acids for | | Leccino | | 112.00 | 131.00 | | |
| Kalahari | | Dritta | * | 314.00 | 435.00 | Cytolase 0 | Ranalli <i>et al</i> . |
| melon seeds | | | | | 427.00 | Maxoliva | (2003) |
| | | | | | 388.00 | Bioliva | |
| | | Caroleo | * | 222.00 | 329.00 | Cytolase 0 | |
| | | | | | 318.00 | Maxoliva | |
| | | | | | 287.00 | Bioliva | |
| | | Coratina | * | 382.00 | 479.00 | Cytolase 0 | |
| | | | | | 462.00 | Maxoliva | |
| | | | | | 431.00 | Bioliva | |
| | | Coratina | * | 691.30 | 751.00 | A / B / C** (1:1:1) | Aliakbarian |
| | | | | | | | et al. (2008) |
| | | Coratina | * | 574.50 | 804.30 | A / B / C** (1:1:1) | De Faveri et |
| | | | | | | | al. (2008) |
| | | Koroneiki | * | 179.00 | 309.00 | Pectinex | Najafian <i>et</i> |
| | | | | | 245.00 | Pectinase | al. (2009) |
| | | Iranian | * | 302.33 | 357.67 | Pectinex | |
| | | oleaginous | | | 359.00 | Pectinase | |
| | | Mission | * | 199.67 | 306.67 | Pectinex | |
| | | | | | 258.33 | Pectinase | |
| Palm fruit | | * | 21.43 | 17.43 | Pectinase / Cellulase | Teixeira et a | |
| | | | | | 14.76 | Pectinase / Cellulase / | (2013) |
| | | | | | | Tannase | |
| | | | | | 26.43 | Tannase | |

The column adjacent to the olive paste refers to the different olive species used.

*data not reported

**A: pectinase, cellulase, hemicellulase; B: pectinase, hemicellulase; C: pectolytic enzyme